=> fil reg; d stat que 18; fil capl; d que nos 19

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VAR G1=O/S
VAR G2=OH/SH/NH2
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES: RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 18

STEREO ATTRIBUTES: NONE

13 SEA FILE=REGISTRY SSS FOLKE OF

100.0% PROCESSED 7857 ITERATIONS

SEARCH TIME: 00.00.03

13 Answers

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FILE COVERS 1967 - 16 Mar 2001 VOL 134 ISS 13 Searched by Barb O'Bryen, STIC 308-4291 ·FILE LAST UPDATED: 15 Mar 2001 (20010315/ED)

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L1 STR L8 13 SEA FILE=REGISTRY SSS FUL L1

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L1 STR L8 13 SEA FILE=REGISTRY SSS FUL L1

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PROCESSING COMPLETED FOR L10
Searched by Barb O'Bryen, STIC 308-4291

L12 23 DUP REMETO 42 DUPLICATES REMOVED ANSWERS '1-22' FROM FILE CAPLUS ANSWER '23' FROM FILE USPATFULL

dibib abs hitstr 112 1-23% fil cao; d que nos 111; fil hom

CAPLUS COPYRIGHT 2001 ACS L12 ANSWER 1 OF 23

DUPLICATE 1

ACCESSION NUMBER:

1997:492901 CAPLUS

DOCUMENT NUMBER:

127:156724

TITLE:

Lipoxin compounds, and preparation thereof, for modulation of inflammation related to columnar

epithelia

INVENTOR (S):

Madara, James L.; Serhan, Charles N.; Colgan, Sean P.

PATENT ASSIGNEE(S):

SOURCE:

U.S., 19 pp. Cont.-in-part of U.S. Ser. No. 84,311,

abandoned. CODEN: USXXAM

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. US 1994-268049 US 5650435 Α 19970722 US 1997-806278 А 20000808 US 6100296 US 2000-496717 US 6177468 в1 20010123

PRIORITY APPLN. INFO.:

US 1991-677388 19910401 US 1991-748349 19910822 US 1993-84311 19930629

19940629

19970225

20000202

US 1994-268049 19940629 US 1997-806278 19970225 US 1997-955860 19971021

Pharmaceutical compns. contg. lipoxin compds. and therapeutic uses for the AΒ compds. in treating or preventing a disease or condition assocd. with columnar epithelial inflammation are provided. Also disclosed are methods for screening for compds. useful in preventing columnar epithelial inflammation. The compds. include lipoxin A4 and analogs thereof. Compds. were tested for their ability to inhibit neutrophil transmigration on epithelial cells. Prepn. of lipoxin compds. is also described.

193279-94-6 193611-38-0 IT

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(lipoxin compds., and prepn. thereof, for modulation of inflammation related to columnar epithelia)

193279-94-6 CAPLUS RN

CN

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, (5S, 6R, 7E, 9E, 11Z, 13E) - [partial] - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

RN 193611-38-0 CAPLUS

CN 7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, methyl ester, (5s,6R,7E,9E,11Z,13E)-[partial]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

MeO
$$(CH_2)_3$$
 S R E E Z OPh OH

L12 ANSWER 2 OF 23 CAPLUS COPYRIGHT 2001 ACS DUPLICATE 2

ACCESSION NUMBER: 1995:795408 CAPLUS

DOCUMENT NUMBER: 124:8499

TITLE: Lipoxin compounds INVENTOR(S): Serhan, Charles N.

PATENT ASSIGNEE(S): Brigham and Women's Hospital, USA

SOURCE: U.S., 4 pp.
CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 4

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE	
	-·				
US 5441951	A	19950815	US 1994-260030	19940615	
US 5648512	·A	19970715	US 1995-453125	19950531	
US 6048897	A	20000411	US 1996-712610	19960913	
PRIORITY APPLN. INF	0.:		US 1993-77300	19930615	
•			US 1994-260030	19940615	
			US 1995-453125	19950531	

OTHER SOURCE(S): MARPAT 124:8499

AB Compds. having the active site of natural lipoxins, but a longer tissue half-life are disclosed. These mols. are useful for treating vasoconstrictive, inflammatory, myeloid suppressive, cardiovascular, and gastrointestinal diseases.

IT 161718-15-6P 161718-22-5P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(lipoxin analogs with longer tissue half-life)

RN 161718-15-6 CAPLUS

CN 7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy- (9CI) (CA INDEX NAME)

PAGE 1-B

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RN 161718-22-5 CAPLUS

CN 7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, methyl ester (9CI) (CA INDEX NAME)

PAGE 1-A

PAGE 1-B

-oPh

L12 ANSWER 3 OF 23 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:666685 CAPLUS

DOCUMENT NUMBER:

133:256820

TITLE:

Lipoxin compounds and their use

Serhan, Charles N.

INVENTOR(S):
PATENT ASSIGNEE(S):

Brigham and Women's Hospital, USA

SOURCE:

PCT Int. Appl., 86 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000055109	A1	20000921	WO 2000-US6583	20000314

W: AU, CA, JP

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE

PRIORITY APPLN. INFO.:

US 1999-125209 19990318

AB Aspirin (ASA) triggers a switch in the biosynthesis of lipid mediators, inhibiting prostanoid prodn. and initiating 15-epi-lipoxin generation, through the acetylation of cyclooxygenase II. Results of expts. indicated that the inhibitory actions of aspirin-triggered lipoxins (ATL) are both tissue- and delivery site-dependent and are the first to show that stable analogs of ATL inhibit acute inflammation at sites distant from the point of delivery. Since ATL stable analogs were designed as mimetics to incorporate the native aspirin-triggered structural features, the findings provide new tools to examine endogenous anti-inflammatory pathways as well as avenues to approach the development of both topical and i.v. anti-PMN therapies.

IT 228549-33-5, ATLa2

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)
Searched by Barb O'Bryen, STIC 308-4291

(aspirin-triggered lipoxins in inflammation inhibition and drug delivery)

RN 228549-33-5 CAPLUS

> 7,9,11,13-Hexadecatetraenoic acid, 16-(4-fluorophenoxy)-5,6,15-trihydroxy-, (5s, 6r, 7e, 9e, 11z, 13e, 15s) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

REFERENCE COUNT:

REFERENCE (S):

- (1) Brigham & Women's Hospital; WO 9811049 A 1998
- (2) Takano, T; 1998, 21, P41 CAPLUS(3) Takano, T; J CLIN INVEST 1998, V101(4), P819 CAPLUS

CAPLUS COPYRIGHT 2001 ACS L12 ANSWER 4 OF 23

ACCESSION NUMBER:

2000:666595 CAPLUS

DOCUMENT NUMBER:

133:247276

TITLE:

CN

Use of lipoxin compounds for inhibiting of

TNF-.alpha.-initiated neutrophil response

INVENTOR (S): Serhan, Charles N.

Brigham and Women's Hospital, USA PATENT ASSIGNEE(S):

SOURCE:

PCT Int. Appl., 93 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000054767	A1	20000921	WO 2000-US6582	20000314

W: AU, CA, JP

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,

PT, SE

PRIORITY APPLN. INFO.:

US 1999-125205 19990318

OTHER SOURCE(S):

MARPAT 133:247276

The impact of lipoxin A4 (LXA4) and aspirin-triggered-lipoxins (ATL) was AB investigated in tumor necrosis factor (TNF.alpha.)-initiated neutrophil (PMN) responses in vitro and in vivo using metabolically stable LX analogs. At concns. as low as 1-10 nM, the LXA4 and ATL analogs each inhibited TNF.alpha.-stimulated superoxide anion generation and IL-1.beta. release by human PMN.

IT 171030-12-9

> RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(lipoxin compds. for inhibiting of TNF-.alpha.-initiated neutrophil response)

RN 171030-12-9 CAPLUS

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, CN (5S, 6R, 7E, 9E, 11Z, 13E, 15R) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

REFERENCE COUNT:

REFERENCE (S):

7

(1) Brigham & Women'S Hospital; WO 9429262 A 1994 CAPLUS

(2) Brigham & Women'S Hospital; WO 9501179 A 1995 CAPLUS

(3) Brigham & Women'S Hospital; WO 9811049 A 1998 CAPLUS

(5) Serhan, C; US 5441951 A 1995 CAPLUS

(6) Takano, T; JOURNAL OF CLINICAL INVESTIGATIONS 1998, V101(4), P819 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 5 OF 23 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

2000:666589 CAPLUS

DOCUMENT NUMBER:

133:232831

TITLE:

Lipoxin analogs for regulation of phospholipase D

activity, and therapeutic use thereof

INVENTOR (S):

Serhan, Charles N.

PATENT ASSIGNEE (S):

Brigham and Women's Hospital, USA

SOURCE:

PCT Int. Appl., 77 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent English

LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
WO 2000054761	A2	20000921	WO 2000-US6669	20000314

W: AU, CA, JP

RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,

PT, SE

PRIORITY APPLN. INFO.:

US 1999-125194 19990318

OTHER SOURCE(S):

MARPAT 133:232831 Lipoxin analogs are provided for modulating diseases and conditions assocd. with phospholipase D activity, including those assocd. with phospholipase D-initiated superoxide generation or degranulation activity. The effect of 15-epi-LXA4 in polyisoprenylphosphate signaling was examd. 15-Epi-LXA4 inhibited LTB4-stimulated phospholipase D activity and superoxide generation.

IT 171030-14-1

RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study)

(lipoxin analogs for regulation of phospholipase D activity, and therapeutic use)

RN 171030-14-1 CAPLUS

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, methyl CN ester, (5S, 6R, 7E, 9E, 11Z, 13E, 15R) - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

MeO
$$(CH_2)_3$$
 S R E E Z E R OPh

IT 205176-29-0

> RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(lipoxin analogs for regulation of phospholipase D activity, and

therapeutic use)

205176-29-0 CAPLUS ·RN ·

7,9,11,13-Hexadecatetraenoic acid, 16-(4-fluorophenoxy)-5,6,15-trihydroxy-CN , methyl ester, (5S,6R,7E,9E,11Z,13E,15S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

PAGE 1-A

PAGE 1-B

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SOURCE:

CORPORATE SOURCE:

L12 ANSWER 6 OF 23 CAPLUS COPYRIGHT 2001 ACS

2000:198492 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 133:3608

TITLE: Lipoxin A4 Analogues Inhibit Leukocyte Recruitment to

Porphyromonas gingivalis: A Role for Cyclooxygenase-2

and Lipoxins in Periodontal Disease

AUTHOR (S): Pouliot, Marc; Clish, Clary B.; Petasis, Nicos A.; Van

Dyke, Thomas E.; Serhan, Charles N.

Center for Experimental Therapeutics and Reperfusion Injury Department of Anesthesiology Perioperative and Pain Medicine Brigham and Women's Hospital, Harvard

Medical School, Boston, MA, 02115, USA

Biochemistry (2000), 39(16), 4761-4768

CODEN: BICHAW; ISSN: 0006-2960

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal

LANGUAGE: English The potential involvement of the inducible cyclooxygenase isoform (COX-2) and the role of novel lipid mediators were investigated in the

pathogenesis of periodontal disease. Crevicular fluids from localized juvenile periodontitis (LJP) patients contained prostaglandin (PG)E2 and 5-lipoxygenase-derived products, leukotriene B4, and the biosynthesis Searched by Barb O'Bryen, STIC 308-4291

interaction product, lipoxin (LX)A4. Neutrophils from peripheral blood of LJP patients, but not from asymptomatic donors, also generated LXA4, suggesting a role for this immunomodulatory mol. in periodontal disease. To characterize host responses of interest to periodontal pathogens, P. gingivalis was introduced within murine dorsal air pouches. In the air pouch cavity, P. gingivalis elicited leukocyte infiltration, concomitant with elevated PGE2 levels in the cellular exudates, and upregulated COX-2 expression in infiltrated leukocytes. In addn., human neutrophils exposed to P. gingivalis also upregulated COX-2 expression. Blood borne P. gingivalis gave increases in the murine tissue levels of COX-2 mRNA assocd. with both heart and lungs, supporting a potential role for this oral pathogen in the evolution of systemic events. The administration of metabolically stable analogs of LX and of aspirin-triggered LX potently blocked neutrophil traffic into the dorsal pouch cavity and lowered PGE2 levels within exudates. These results identify thus PMN as an addnl. and important source of PGE2 in periodontal tissues. Moreover, they provide evidence for a novel protective role for LX in periodontitis, limiting further PMN recruitment and PMN-mediated tissue injury that can lead to loss of inflammatory barriers that prevent systemic tissue invasion of oral microbial pathogens.

IT 230954-56-0

> RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (lipoxin A4 analogs inhibit leukocyte recruitment to Porphyromonas gingivalis in periodontal disease)

RN 230954-56-0 CAPLUS

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, CN (5S, 6R, 7E, 9E, 11Z, 13E, 15S) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

REFERENCE COUNT:

REFERENCE(S):

- (1) Abramson, M; J Periodont Res 1992, V27, P539 CAPLUS
- (2) Albers, H; Dtsch Zahnarztl Z 1979, V34, P440 CAPLUS
- (3) Assuma, R; J Immunol 1998, V160, P403 CAPLUS(4) Babior, B; Blood 1984, V64, P959 CAPLUS
- (5) Borgeat, P; Clin Biochem 1990, V23, P459 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 7 OF 23 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: DOCUMENT NUMBER:

2000:146214 CAPLUS

132:303164

TITLE:

Cutting edge: lipoxin (LX) A4 and aspirin-triggered

15-Epi-LXA4 block allergen-induced eosinophil

trafficking

AUTHOR (S):

Bandeira-Melo, Christianne; Bozza, Patricia T.; Diaz,

Bruno L.; Cordeiro, Renato S. B.; Jose, Peter J.;

Martins, Marco A.; Serhan, Charles N.

CORPORATE SOURCE:

Department of Physiology and Pharmacodynamics, Oswaldo

Cruz Institute, Rio de Janeiro, Brazil

SOURCE:

J. Immunol. (2000), 164(5), 2267-2271 Searched by Barb O'Bryen, STIC 308-4291

CODEN: JOIMA3; ISSN: 0022-1767

PUBLISHER:

The street of the

American Association of Immunologists

DOCUMENT TYPE:

English

LANGUAGE: AB

Tissue eosinophilia prevention represents one of the primary targets to new anti-allergic therapies. As lipoxin A4 (LXA4) and aspirin-triggered 15-epi-LXA4 (ATL) are emerging as endogenous "stop signals" produced in distinct pathologies including some eosinophil-related pulmonary disorders, the authors evaluated the impact of in situ LXA4/ATL

metabolically stable analogs on allergen-induced eosinophilic pleurisy in sensitized rats. LXA4/ATL analogs dramatically blocked allergic pleural eosinophil influx, while concurrently increasing circulating eosinophilia, inhibiting the earlier edema and neutrophilia assocd. with allergic The mechanisms underlying this LXA4/ATL-driven allergic reaction. eosinophilia blockade was independent of mast cell degranulation and involved LXA4/ATL inhibition of both IL-5 and eotaxin generation, as well as platelet activating factor action. These findings reveal LXA4/ATL as a novel class of endogenous anti-allergic mediators, capable of preventing local eosinophilia.

ΙT 265316-15-2

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(antiallergic impact of lipoxin (LX) A4 and aspirin-triggered

15-Epi-LXA4 analogs in eosinophilic pleurisy)

RN 265316-15-2 CAPLUS

7,9,11,13-Eicosatetraenoic acid, 16-(4-fluorophenoxy)-5,6,15-trihydroxy-, CN (5S, 6R, 7E, 9E, 11Z, 13E, 15S) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

REFERENCE COUNT: REFERENCE(S):

36

(1) Alves, A; Eur J Pharmacol 1996, V312, P89 CAPLUS

(4) Clish, C; Proc Natl Acad Sci USA 1999, V96, P8247 CAPLUS

(7) Edenius, C; FEBS Lett 1990, V272, P25 CAPLUS

(8) Gewirtz, A; Am J Physiol 1999, V276, PC988 CAPLUS

(9) Gewirtz, A; J Clin Invest 1998, V101, P1860 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2001 ACS L12 ANSWER 8 OF 23

ACCESSION NUMBER:

2000:138534 CAPLUS

DOCUMENT NUMBER:

132:278030

TITLE:

Cutting edge: lipoxins rapidly stimulate

non-phlogistic phagocytosis of apoptotic neutrophils

by monocyte-derived macrophages

Godson, Catherine; Mitchell, Siobhan; Harvey, Killeen;

Petasis, Nicos A.; Hogg, Nancy; Brady, Hugh R.

Centre for Molecular Inflammation and Vascular

Research, Mater Misericordiae Hospital and Department

of Medicine and Therapeutics, Conway Institute of Biomolecular and Biomedical Research, University

College Dublin, Dublin, 7, Ire. Searched by Barb O'Bryen, STIC 308-4291

AUTHOR (S):

CORPORATE SOURCE:

SOURCE:

J. Immunol. (2000), 164(4), 1663-1667

CODEN: JOIMA3; ISSN: 0022-1767

PUBLISHER:

American Association of Immunologists

DOCUMENT TYPE:

Journal English

LANGUAGE:

AB

Lipoxins (LX) are lipoxygenase-derived eicosanoids generated during inflammation. LX inhibit polymorphonuclear neutrophil (PMN) chemotaxis and adhesion and are putative braking signals for PMN-mediated tissue injury. Here, the authors report that LXA4 promotes another important step in the resoln. phase of inflammation, namely, phagocytosis of apoptotic PMN by monocyte-derived macrophages (M.phi.). LXA4 triggered rapid, concn.-dependent uptake of apoptotic PMN. This bioactivity was shared by stable synthetic LXA4 analogs but not by other eicosanoids LXA4-triggered phagocytosis did not provoke IL-8 or monocyte chemoattractant protein-1 release. LXA4-induced phagocytosis was attenuated by anti-CD36, .alpha.v.beta.3, and CD18 mAbs. LXA4-triggered PMN uptake was inhibited by pertussis toxin and by 8-bromo-cAMP and was mimicked by Rp-cAMP, a protein kinase A inhibitor. LXA4 attenuated PGE2-stimulated protein kinase A activation in M.phi.. These results suggest that LXA4 is an endogenous stimulus for PMN clearance during

inflammation and provide a novel rationale for using stable synthetic

ΙT 260064-29-7

> RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(lipoxins rapidly stimulate non-phlogistic phagocytosis of apoptotic neutrophils by monocyte-derived macrophages)

260064-29-7 CAPLUS RN

7,9,11,13-Eicosatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, CN (5S, 6R, 7E, 9E, 11Z, 13E, 15R) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

analogs as anti-inflammatory compds. in vivo.

REFERENCE COUNT:

REFERENCE (S):

(1) Asch, A; Science 1993, V262, P1436 CAPLUS

(2) Chiang, N; J Pharmacol Exp Ther 1998, V287, P779 CAPLUS

(3) Clish, C; Proc Natl Acad Sci USA 1999, V96, P8247 CAPLUS

(4) Colgan, S; J Clin Invest 1993, V92, P75 CAPLUS

(5) Devitt, A; Nature 1998, V392, P505 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

COPYRIGHT 2001 ACS L12 ANSWER 9 OF 23 CAPLUS

ACCESSION NUMBER:

2000:228388 CAPLUS

DOCUMENT NUMBER:

133:16222

TITLE:

AUTHOR (S):

Activation of lipoxin A4 receptors by

aspirin-triggered lipoxins and select peptides evokes

ligand-specific responses in inflammation

Chiang, Nan; Fierro, Iolanda M.; Gronert, Karsten;

Serhan, Charles N.

CORPORATE SOURCE:

Center for Experimental Therapeutics and Reperfusion Injury, Department of Anesthesiology, Pe Searched by Barb O'Bryen, STIC 308-4291 Perioperative

and Pain Medicine, Brigham and Women's Hospital and Harvard Medical School, Boston, MA, 02115, USA

J. Exp. Med. (2000), 191(7), 1197-1207

CODEN: JEMEAV; ISSN: 0022-1007 Rockefeller University Press

PUBLISHER: DOCUMENT TYPE:

SOURCE:

Journal English

LANGUAGE: Lipoxin (LX) A4 and aspirin-triggered LX (ATL) are endogenous lipids that regulate leukocyte trafficking via specific LXA4 receptors (ALXRs) and mediate antiinflammation and resoln. ATL analogs dramatically inhibited human neutrophil [polymorphonuclear leukocyte (PMN)] responses evoked by a potent necrotactic peptide derived from mitochondria as well as a rogue synthetic chemotactic peptide. These bioactive lipid analogs and small peptides each selectively competed for specific 3H-LXA4 binding with recombinant human ALXR, and its N-glycosylation proved essential for peptide but not LXA4 recognition. Chimeric receptors constructed from receptors with opposing functions, namely ALXR and leukotriene B4 receptors (BLTs), revealed that the seventh transmembrane segment and adjacent regions of ALXR are essential for LXA4 recognition, and addnl. regions of ALXR are required for high affinity binding of the peptide ligands. Together, these findings are the first to indicate that a single 7-transmembrane receptor can switch recognition as well as function with certain chemotactic peptides to inhibitory with ATL and LX (lipid ligands). Moreover, they suggest that ALXR activation by LX or ATL can protect the host from potentially deleterious PMN responses assocd. with

recognition of peptide fragments. IT 205176-29-0

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311.

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RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study)

(lipoxin A4 receptors activation by aspirin-triggered lipoxins and select peptides evokes ligand-specific responses in inflammation)

innate immunity as well as direct effector responses in tissue injury by

RN 205176-29-0 CAPLUS

CN 7,9,11,13-Hexadecatetraenoic acid, 16-(4-fluorophenoxy)-5,6,15-trihydroxy-, methyl ester, (5S,6R,7E,9E,11Z,13E,15S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

PAGE 1-B

__OMe

REFERENCE COUNT: REFERENCE(S):

48

(1) Beckman, E; Nature 1994, V372, P691 CAPLUS

(2) Brezinski, D; Biol Mass Spectrom 1991, V20, P45 CAPLUS

(3) Castano, A; Science 1995, V269, P223 CAPLUS Searched by Barb O'Bryen, STIC 308-4291

(4) Chiang, N; J Clin Invest 1999, V104, P309 CAPLUS

(5) Clish, C; Proc Natl Acad Sci USA 1999, V96, P8247

CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2001 ACS L12 ANSWER 10 OF 23

ACCESSION NUMBER:

2000:354308 CAPLUS

DOCUMENT NUMBER:

134:4031

TITLE:

A novel polyisoprenyl phosphate signaling cascade in

human neutrophils

AUTHOR (S):

Levy, Bruce D.; Serhan, Charles N.

CORPORATE SOURCE:

Center for Experimental Therapeutics and Reperfusion Injury, Department of Anesthesia, Perioperative and Pain Medicine, Brigham and Women's Hospital and Harvard Medical School, Boston, MA, 02115, USA

SOURCE:

Ann. N. Y. Acad. Sci. (2000), 905 (Lysophospholipids

and Eicosanoids in Biology and Pathophysiology), 69-80

CODEN: ANYAA9; ISSN: 0077-8923

PUBLISHER:

New York Academy of Sciences

DOCUMENT TYPE:

Journal

LANGUAGE: English Activation of neutrophil (PMN) surface receptors can evoke inflammation and tissue injury via aberrant release of excess effectors. The mol. mechanisms involved in host protection and control of PMN responses have yet to be defined. As Billah and coworkers (1989) and Exton (1997), for example, have pointed out, phospholipase D (PLD) signaling is known to play a pivotal role in PMN activation. Here, we detd. the relationship between polyisoprenyl phosphate (PIPP) remodeling and PLD signaling and their impact in activation of PMN receptors by "proinflammatory" (leukotriene B4), and "anti-inflammatory" (aspirin-triggered lipoxin A4) ligands. Activation of the leukotriene B4 receptor initiated a rapid (within seconds) decrement in presqualene diphosphate (PSDP), activation of PLD and prodn. of superoxide anions. This contrasts with activation of the LXA4 receptor by an aspirin-triggered lipoxin A4 mimetic that before leukotriene B4 gave an inverse relationship with rapidly increasing PSDP levels, and inhibition of both PLD activity and superoxide generation. PSDP proved to be a potent and direct-acting inhibitor of PLD (rhPLD1b: Ki = 5.9 nM), a property not shared by structurally related endogenous This PIPP also interacted with Src homol. domains, selectively targeting SH2 and not SH3 domains. These results indicate a role for ligand-driven rapid PIPP remodeling as an early switch and "stop" signaling event that controls PMN. Moreover, they indicate that PSDP directly down-regulates PMN signaling events via select protein-targeted interactions controlling intracellular responses relevant in inflammation.

IT 205176-29-0

RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study)

(polyisoprenyl phosphate remodeling and phospholipase D signaling in relation to activation of leukotriene B4 receptor and lipoxin A4 receptor in neutrophils)

205176-29-0 CAPLUS

7,9,11,13-Hexadecatetraenoic acid, 16-(4-fluorophenoxy)-5,6,15-trihydroxy-CN , methyl ester, (5S, 6R, 7E, 9E, 11Z, 13E, 15S) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

PAGE 1-A

PAGE 1-B

- OMe

REFERENCE COUNT:

REFERENCE(S):

(1) Bach, T; Lipids 1995, V30, P191 CAPLUS

(2) Billah, M; J Biol Chem 1989, V264, P17069 CAPLUS

(3) Chardin, P; FEBS Lett 1995, V369, P47 CAPLUS

(4) Chiang, N; J Pharmacol Exp Ther 1998, V287, P779

CAPLUS

(5) Exton, J; J Biol Chem 1997, V272, P15579 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 11 OF 23

ACCESSION NUMBER:

CAPLUS COPYRIGHT 2001 ACS 1999:507837 CAPLUS

DOCUMENT NUMBER:

131:266744

TITLE:

Local and systemic delivery of a stable

aspirin-triggered lipoxin prevents neutrophil-

recruitment in vivo

AUTHOR (S):

SOURCE:

Clish, Clary B.; O'Brien, Jennifer A.; Gronert,

Karsten; Stahl, Gregory L.; Petasis, Nicos A.; Serhan,

Charles N.

CORPORATE SOURCE:

Center for Experimental Therapeutics and Reperfusion Injury, Department of Anesthesiology, Perioperative and Pain Medicine, Brigham and Women's Hospital and

Harvard Medical School, Boston, MA, 02115, USA Proc. Natl. Acad. Sci. U. S. A. (1999), 96(14),

8247-8252

CODEN: PNASA6; ISSN: 0027-8424 National Academy of Sciences

DOCUMENT TYPE:

PUBLISHER:

Journal

English LANGUAGE:

Aspirin (ASA) triggers a switch in the biosynthesis of lipid mediators, inhibiting prostanoid prodn. and initiating 15-epi-lipoxin generation through the acetylation of cyclooxygenase II. These aspirin-triggered lipoxins (ATL) may mediate some of ASA's beneficial actions and therefore are of interest in the search for novel antiinflammatories that could manifest fewer unwanted side effects. Here, we report that design modifications to native ATL structure prolong its biostability in vivo. In mouse whole blood, ATL analogs protected at carbon 15 [15(R/S)-methyl-lipoxin A4 (ATLal)] and the omega end [15-epi-16-(parafluoro)-phenoxy-LXA4 (ATLa2)] were recoverable to .apprxeq.90 and 100% at 3 h, resp., compared with a .apprxeq.40% loss of native lipoxin A4. retains bioactivity and, at levels as low as .apprxeq.24 nmol/mouse, potently inhibited tumor necrosis factor-.alpha.-induced leukocyte recruitment into the dorsal air pouch. Inhibition was evident by either local intra-air pouch delivery (.apprxeq.77% inhibition) or systemic Searched by Barb O'Bryen, STIC 308-4291

delivery by i.v. injection (.apprxeq.85% inhibition) and proved more potent than local delivery of ASA. Rank order for inhibiting polymorphonuclear leukocyte infiltration was: ATLa2 (10 .mu.g, i.v.) .apprxeq.ATLa2 (10 .mu.g, local) .apprxeq.dexamethasone (10 .mu.g, local) >ASA (1.0 mg, local). Applied topically to mouse ear skin, ATLa2 also inhibited polymorphonuclear leukocyte infiltration induced by leukotriene B4 (.apprxeq.78% inhibition) or phorbol ester (.apprxeq.49% inhibition), which initiates endogenous chemokine prodn. These results indicate that this fluorinated analog of natural aspirin-triggered lipoxin A4 is bioavailable by either local or systemic delivery routes and is a more potent and precise inhibitor of neutrophil accumulation than is ASA.

IT 228549-33-5

RN

CN

RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); THU (Therapeutic use); BIOL (Biological study); PROC (Process); USES (Uses)

(structure on p. 8249; local and systemic delivery of a stable aspirin-triggered lipoxin prevents neutrophil recruitment in vivo) 228549-33-5 CAPLUS

7,9,11,13-Hexadecatetraenoic acid, 16-(4-fluorophenoxy)-5,6,15-trihydroxy-, (5S, 6R, 7E, 9E, 11Z, 13E, 15S) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

REFERENCE COUNT:

REFERENCE(S):

31

- (2) Bradley, P; J Invest Dermatol 1982, V78, P206 CAPLUS
- (3) Chavis, C; Biochem Biophys Res Commun 1995, V207, P273 CAPLUS
- (4) Chavis, C; J Exp Med 1996, V183, P1633 CAPLUS
- (5) Chiang, N; J Pharmacol Exp Ther 1998, V287, P779 CAPLUS
- (6) Dahlen, S; Lipoxygenases and Their Products 1991, P235 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

CAPLUS COPYRIGHT 2001 ACS L12 ANSWER 12 OF 23

ACCESSION NUMBER:

1999:797035 CAPLUS

DOCUMENT NUMBER:

132:189458

TITLE: ·

Anti-inflammatory actions of lipoxin A4 stable analogs are demonstrable in human whole blood: modulation of

leukocyte adhesion molecules and inhibition of

neutrophil-endothelial interactions

AUTHOR (S):

Filep, Janos G.; Zouki, Christine; Petasis, Nicos A.;

CORPORATE SOURCE:

Hachicha, Mohamed; Serhan, Charles N. Research Center, Maisonneuve-Rosemont Hospital,

Department of Medicine, University of Montreal,

Montreal, PQ, Can.

SOURCE:

Blood (1999), 94(12), 4132-4142 CODEN: BLOOAW; ISSN: 0006-4971

W. B. Saunders Co.

DOCUMENT TYPE:

Journal English

LANGUAGE:

PUBLISHER:

The authors examd. in whole blood the actions of 2 lipoxin A4 (LXA4) stable analogs, 15-R/S-methyl-LXA4 and 16-phenoxy-LXA4, for their impact on the expression of adhesion mols. on human leukocytes and coronary artery endothelial cells (HCAEC) and on neutrophil adhesion to HCAEC in Both LXA4 analogs in nanomolar to micromolar concns. prevented shedding of L-selectin and downregulated CD11/CD18 expression on resting neutrophils, monocytes, and lymphocytes. Changes in CD11/CD18 expression were blocked by the mitogen-activated protein kinase kinase inhibitor The LXA4 analogs also attenuated changes in L-selectin and CD11/CD18 expression evoked by platelet-activating factor (PAF), interleukin-8, or C-reactive protein-derived peptide 201-206 with IC50 values of 0.2 to 1.9 .mu.mol/L, whereas they did not affect lipopolysaccharide (LPS) - or tumor necrosis factor -. alpha. - stimulated expression of E-selectin and intercellular adhesion mol.-1 on HCAEC. These LXA4 analogs markedly diminished adhesion of neutrophils to LPS-activated HCAEC. Inhibition of adhesion was additive with function blocking anti-E-selectin and anti-L-selectin antibodies, but was not additive with anti-CD18 antibody. Combining LXA4 analogs with dexamethasone (100 nmol/L) almost completely inhibited PAF-induced changes in adhesion mol. expression on leukocytes and gave additive inhibition of neutrophil adhesion to HCAEC. Culture of HCAEC with dexamethasone, but not with LXA4 analogs, also decreased neutrophil attachment. Thus, LXA4 stable analogs modulate expression of both L-selectin and CD11/CD18 on resting and immunostimulated leukocytes and inhibit neutrophil adhesion to HCAEC by attenuating CD11/CD18 expression. These actions are additive with those of glucocorticoids and may represent a novel and potent regulatory mechanism by which LXA4 and aspirin-triggered 15-epi-LXA4 modulate leukocyte trafficking.

260064-29-7

IT

: AB

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (anti-inflammatory actions of lipoxin A4 stable analogs in human whole blood are via modulation of leukocyte adhesion mols. and inhibition of neutrophil-endothelial interactions)

RN 260064-29-7 CAPLUS

CN 7,9,11,13-Eicosatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, (5s,6R,7E,9E,11Z,13E,15R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

$$\begin{array}{c|c}
\text{OPh} & \text{OH} \\
\text{N-Bu} & \text{E} & \text{E} \\
\text{OH} & \text{OH}
\end{array}$$

REFERENCE COUNT:

REFERENCE (S).:

39

- (1) Bator, J; Immunopharmacology 1992, V23, P139 CAPLUS
- (3) Capodici, C; J Clin Invest 1998, V102, P165 CAPLUS
- (4) Chen, A; J Exp Med 1995, V182, P519 CAPLUS
- (5) Claria, J; Proc Natl Acad Sci USA 1995, V92, P9475 CAPLUS
- (6) Colgan, S; J Clin Invest 1993, V92, P75 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 13 OF 23 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1999:274601 CAPLUS

. DOCUMENT NUMBER:

TITLE:

LXA4, aspirin-triggered 15-epi-LXA4, and their analogs selectively downregulate PMN azurophilic degranulation Gewirtz, Andrew T.; Fokin, Valery V.; Petasis, Nicos

AUTHOR(S): Gewirtz, Andrew T.; Fokin, Valery V.; Pe A.; Serhan, Charles N.; Madara, James L.

CORPORATE SOURCE:

Department of Pathology and Laboratory Medicine, Emory

University School of Medicine, Atlanta, GA, USA Am. J. Physiol. (1999), 276(4, Pt. 1), C988-C994

CODEN: AJPHAP; ISSN: 0002-9513

PUBLISHER:

SOURCE:

American Physiological Society

DOCUMENT TYPE:

Journal

LANGUAGE:

English

The eicosanoid lipoxin A4 (LXA4) is biosynthesized in vivo by cells AB present at inflammatory sites and appears to be an endogenous anti-inflammatory mediator. Further, in the presence of aspirin, the 15-epimer of LXA4 (15-epi-LXA4) is biosynthesized and may mediate some of aspirin's desirable bioactions. LXA4, 15-epi-LXA4, and their stable analogs inhibit inflammation in established animal models, indicating that these compds. may be useful for treating inflammatory disease states. investigate the cellular mechanisms by which these lipid mediators downregulate inflammation, the authors investigated whether these eicosanoids could influence receptor-mediated degranulation of human neutrophils, an event thought to play a major causative role in several inflammatory disease states. LXA4, 15-epi-LXA4, and their stable analogs potently (IC50 <1 nM) and selectively downregulated neutrophil release of azurophilic granule contents but did not affect other neutrophil secretory functions. Thus, the cellular basis of action of these natural off-switches to inflammation appears to involve downregulation of neutrophil azurophilic granule release.

IT 171030-12-9 228549-33-5

RL: BAC (Biological activity or effector, except adverse); BIOL (Biological study)

(LXA4 and aspirin-triggered 15-epi-LXA4 and their analogs selectively downregulate PMN azurophilic degranulation)

RN 171030-12-9 CAPLUS

CN 7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, (5S,6R,7E,9E,11Z,13E,15R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

RN 228549-33-5 CAPLUS CN 7,9,11,13-Hexadecatetraenoic acid, 16-(4-fluorophenoxy)-5,6,15-trihydroxy-, (5S,6R,7E,9E,11Z,13E,15S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

REFERENCE COUNT:

REFERENCE(S):

37

(1) Brunkhorst, B; J Biol Chem 1992, V267, P20659
CAPLUS

(2) Chertov, O; J Exp Med 1997, V186, P739 CAPLUS

(3) Claria, J; Proc Natl Acad Sci USA 1995, V92, P9475 CAPLUS

(4) Clarkson, S; J Exp Med 1986, V164, P474 CAPLUS(5) Cohen, H; J Clin Invest 1978, V61, P1081 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 14 OF 23 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1999:315716 CAPLUS

DOCUMENT NUMBER:

131:101234

TITLE:

Polyisoprenyl phosphate (PIPP) signaling regulates

phospholipase D activity: a "stop" signaling switch

for aspirin-triggered lipoxin A4

AUTHOR (S):

Levy, Bruce D.; Fokin, Valery V.; Clark, Joanna M.; Wakelam, Michael J. O.; Petasis, Nicos A.; Serhan,

Charles N.

CORPORATE SOURCE:

Center for Experimental Therapeutics and Reperfusion

Injury, Brigham and Women's Hospital and Harvard

Medical School, Boston, MA, 02115, USA

SOURCE:

FASEB J. (1999), 13(8), 903-911 CODEN: FAJOEC; ISSN: 0892-6638

PUBLISHER:

Federation of American Societies for Experimental

Biology

DOCUMENT TYPE:

TYPE: Journal

LANGUAGE: English It is of wide interest to understand how opposing extracellular signals AB (pos. or neg.) are translated into intracellular signaling events. Receptor-ligand interactions initiate the generation of bioactive lipids by human neutrophils (PMN), which serve as signals to orchestrate cellular responses important in host defense and inflammation. The authors recently identified a novel polyisoprenyl phosphate (PIPP) signaling pathway and found that one of its components, presqualene diphosphate (PSDP), is a potent neg. intracellular signal in PMN that regulates superoxide anion generation by several stimuli, including phosphatidic The authors detd. intracellular PIPP signaling by autocoids with opposing actions on PMN: leukotriene B4 (LTB4), a potent chemoattractant, and lipoxin A4 (LXA4), a "stop signal" for recruitment. LTB4 receptor activation initiated a rapid decrease in PSDP levels concurrent with activation of PLD and cellular responses. In sharp contrast, activation of the LXA4 receptor reversed LTB4-initiated PSDP remodeling, leading to an accumulation of PSDP and potent inhibition of both PLD and superoxide anion generation. Thus, an inverse relation was established for PSDP levels and PLD activity with two PMN ligands that evoke opposing responses. In addn., PSDP directly inhibited both isolated human recombinant (Ki = 6 nM) and plant (Ki = 20 nM) PLD. Together, these findings link PIPP remodeling to intracellular regulation of PMN function and suggest a role for PIPPs as lipid repressors in signal transduction, a novel mechanism that may also explain aspirin's suppressive actions in vivo in cell signaling.
Searched by Barb O'Bryen, STIC 308-4291

IT 230954-55-9 230954-56-0

RL: BUU (Biological use, unclassified); BIOL (Biological study); USES (Uses)

(increase in intracellular presqualene diphosphate levels and inhibition of phospholipase D activity in superoxide prodn. by human neutrophils by)

RN 230954-55-9 CAPLUS

CN 7,9,11,13-Hexadecatetraenoic acid, 16-(4-fluorophenoxy)-5,6,15-trihydroxy-, methyl ester, (5S,6R,7E,9E,11Z,13E,15R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

PAGE 1-B

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RN 230954-56-0 CAPLUS

CN 7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, (5S,6R,7E,9E,11Z,13E,15S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Double bond geometry as shown.

REFERENCE COUNT: REFERENCE(S):

35

- (1) Abousalham, A; Biochim Biophys Acta 1993, V1158, P1 CAPLUS
- (2) Agwu, D; J Clin Invest 1991, V88, P531 CAPLUS

(3) Bach, T; Lipids 1995, V30, P191 CAPLUS

- (4) Billah, M; J Biol Chem 1989, V264, P17069 CAPLUS
- (6) Chiang, N; J Pharmacol Exp Ther 1998, V287, P779 CAPLUS

ALL CITATIONS AVAILABLE IN THE RE FORMAT

L12 ANSWER 15 OF 23 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1998:180829 CAPLUS

DOCUMENT NUMBER:

128:252973

TITLE:

Lipoxin compounds and their use in treating cell

proliferative disorders

INVENTOR(S):

Serhan, Charles N. Searched by Barb O'Bryen, STIC 308-4291 PATENT ASSIGNEE(S):

Brigham & Women's Hospital, USA

SOURCE:

GI

PCT Int. Appl., 109 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PA	TENT NO.		KI	ND	DATE			A	PPLI	CATIO	ON NO). 	DATE				
	WO	9811049		A.	1	1998	0319		W	0 19	97-US	31634	12	1997	0915			
	•	W: AU, RW: AT,	•	JP CH,	DE,	DK,	ES,	FI,	FR,	GB,	GR,	IE,	IT,	LU,	MC,	NL,	PT,	SE
	US	6048897		A		2000	0411		U	s 19	96-73	12610)	1996	0913			
	AU	9742710		A:	1	1998	0402	*	A	U 19	97-42	2710		1997	0915			
		723321		· B2	2	2000	0824				•			•				
	EP	927150		A:	1	1999	0707		E	P 19	97-94	41078	3	1997	0915			
		R: AT,	BE,	CH,	DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,	
٠.		IE,	FI	• •														
	JP	20015008	66	T	2	2001	0123		J	P 19	98-53	13962	2	1997	0915			
PRI	ORTT	Y APPLN.	INFO	. :					U	s 19	96-73	12610)	1996	0913			
									U	s 19	93-7	7300		1993	0615			
									U	s 19	94-2	6003)	1994	0615			
	•								U	s 19	95-4	53125	5	1995	0531			
									W	0 19	97-U	s163	42	1997	0915			

Lipoxin analogs, e.g., I, having the active site of natural lipoxins but a AΒ longer tissue half-life, are prepd. Thus, I was prepd. via reaction of 3-methyl-3-(trimethylsilyloxy)-1-bromo-1-octene with (7E, 9E, 5S, 6R)-Me 5,6-bis(tert-butyldimethylsilyloxy)-7,9-dodecadien-11-ynoate in benzene contg. propylamine and Pd(PPh3)4 and treating the product with BuN4Fin THF. In particular, 15-epi-lipoxins and their use in ameliorating undesired cell proliferation, which characterizes diseases such as cancer, are also disclosed. The prepd. compds. inhibited neutrophil adhesion to endothelial cells and their transmigration on epithelial cells. Among the prepd. compds., those with acetylenic groups were more stable than others; also the compd. that lacked a 15-OH group showed no biol. activity. [14,15-3H]LXA4 was prepd. and its specific binding to promyelocytic cells (HL-60) was compared with that of [14,15-3H]LTB4 and the results are essentially in agreement with values recently reported by Harada (1991). Bioassays of prepd. compds. were also carried out.

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IT 193611-38-0P

RL: BAC (Biological activity or effector, except adverse); PRP (Properties); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(prepn. of lipoxin analogs for treating cell proliferative disorders)

RN 193611-38-0 CAPLUS

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, methyl CN ester, (5S, 6R, 7E, 9E, 11Z, 13E) - [partial] - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

L12 ANSWER 16 OF 23 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1998:320841 CAPLUS

DOCUMENT NUMBER:

129:62611

TITLE:

Pathogen-induced chemokine secretion from model intestinal epithelium is inhibited by lipoxin A4

analogs

AUTHOR (S):

Gewirtz, Andrew T.; Mccormick, Beth; Neish, Andrew S.; Petasis, Nicos A.; Gronert, Karsten; Serhan, Charles

N.; Madara, James L.

CORPORATE SOURCE:

Department of Pathology and Laboratory Medicine, Emory

University, Atlanta, GA, 30322, USA

SOURCE:

J. Clin. Invest. (1998), 101(9), 1860-1869

CODEN: JCINAO; ISSN: 0021-9738 Rockefeller University Press

PUBLISHER: DOCUMENT TYPE:

Journal

English LANGUAGE:

Enteric pathogens induce intestinal epithelium to secrete chemokines that AB direct movement of polymorphonuclear leukocytes. Mechanisms that might downregulate secretion of these proinflammatory chemokines and thus contain intestinal inflammation have not yet been elucidated. antiinflammatory activities exhibited by the arachidonate metabolite lipoxin A4 (LXA4) suggests that this eicosanoid, which is biosynthesized in vivo at sites of inflammation, might play such a role. We investigated whether chemokine secretion could be regulated by stable analogs of LXA4. Monolayers of T84 intestinal epithelial cells were infected with Salmonella typhimurium, which elicits secretion of distinct apical (pathogen-elicited epithelial chemoattractant) and basolateral (IL-8) chemokines. Stable analogs of LXA4 inhibited S. typhimurium-induced (but not phorbol ester-induced) secretion of both IL-8 and pathogen-elicited epithelial chemoattractant. LXA4 stable analogs did not alter bacterial adherence to nor internalization by epithelia, indicating that LXA4 stable analogs did not block all signals that Salmonella typhimurium activates in intestinal epithelia, but likely led to attenuation of signals that mediate chemokine secretion. Inhibition of S. typhimurium-induced IL-8 secretion by LXA4 analogs was concn.- (IC50.apprx.1 nM) and time-dependent (maximal inhibition .apprx. 1 h). As a result of these effects, LXA4 stable analogs inhibited the ability of bacteria-infected epithelia to direct polymorphonuclear leukocyte movement. These data suggest that LXA4 and its stable analogs may be useful in downregulating active inflammation at mucosal surfaces.

IT 171030-12-9

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RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(pathogen-induced chemokine secretion from model intestinal epithelium is inhibited by lipoxin A4 analogs)

171030-12-9 CAPLUS RN

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, Searched by Barb O'Bryen, STIC 308-4291

(5s, 6r, 7e, 9e, 11z, 13e, 15r) - (9cI)(CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

CAPLUS COPYRIGHT 2001 ACS L12 ANSWER 17 OF 23

ACCESSION NUMBER:

1998:121846

DOCUMENT NUMBER:

128:252676

TITLE:

Neutrophil-mediated changes in vascular permeability

are inhibited by topical application of

aspirin-triggered 15-epi-lipoxin A4 and novel lipoxin

B4 stable analogs

AUTHOR (S):

Takano, Tomoko; Clish, Clary B.; Gronert, Karsten;

Petasis, Nicos; Serhan, Charles N.

CORPORATE SOURCE:

Center for Experimental Therapeutics and Reperfusion Injury, Department of Anesthesia, Harvard Medical School, Brigham and Women's Hospital, Boston, MA,

02115, USA

SOURCE:

J. Clin. Invest. (1998), 101(4), 819-826

CODEN: JCINAO; ISSN: 0021-9738 Rockefeller University Press

DOCUMENT TYPE:

Journal

PUBLISHER: LANGUAGE:

English Neutrophil (PMN) activation is crit. in inflammation and reperfusion injury, suggesting that PMN-directed therapies may be of clin. use. Here, leukotriene B4 (LTB4)-induced PMN influx in ear skin was equiv. between 5-lipoxygenase knockout and wild-type mice. To explore actions of lipoxin (LX) in PMN-mediated tissue injury, we prepd. several novel LX stable analogs, including analogs of LXA4 and aspirin-triggered 15-epi-LXA4 as well as LXB4, and examd. their impact in PMN infiltration and vascular permeability. Each applied topically to mouse ears inhibited dramatically PMN-mediated increases in vascular permeability (IC50 range of 13-26 nmol) with a rank order of 15(R/S)-methyl-LXA4 > 16-para-fluoro-phenoxy-LXA4 .apprx. 5(S)-methyl-LXB4 .gtoreq. 16-phenoxy-LXA4 > 5(R)-methyl-LXB4. These LX mimetics were as potent as an LTB4 receptor antagonist, yet results from microphysiometry with mouse leukocytes indicated that they do not act as LTB4 receptor level antagonists. In addn., within 24 h of delivery, > 90% were cleared from ear biopsies. Neither IL-8, FMLP, C5a, LTD4, nor platelet-activating factor act topically to promote PMN influx. When applied with LTB4, PGE2 enhanced sharply both infiltration and vascular permeability, which were inhibited by a fluorinated stable analog of aspirin-triggered LX. These results indicate that mimetics of LXs and aspirin-triggered 15-epi-LXA4 are topically active in this model and are potent inhibitors of both PMN infiltration and PMN-mediated vascular

IT 171030-14-1 205176-29-0

injury.

RL: BAC (Biological activity or effector, except adverse); THU

(Therapeutic use); BIOL (Biological study); USES (Uses)

(neutrophil-mediated changes in vascular permeability inhibited by topical application of aspirin-triggered 15-epi-lipoxin A4 and novel lipoxin B4 analogs)

171030-14-1 CAPLUS

RN 7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, methyl Searched by Barb O'Bryen, STIC 308-4291 CN

ester, (5S, 6R, 7E, 9E, 11Z, 13E, 15R) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

MeO
$$(CH_2)_3$$
 S E E E E C CH_2 OPh OH

CAPLUS 205176-29-0

7,9,11,13-Hexadecatetraenoic acid, 16-(4-fluorophenoxy)-5,6,15-trihydroxy-CN , methyl ester, (5S,6R,7E,9E,11Z,13E,15S)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

PAGE 1-B

- OMe

CAPLUS COPYRIGHT 2001 ACS L12 ANSWER 18 OF 23

ACCESSION NUMBER:

1997:581522 CAPLUS

DOCUMENT NUMBER:

127:257275

TITLE:

Lipoxin A4 stable analogs inhibit leukocyte rolling

and adherence in the rat mesenteric microvasculature:

role of P-selectin

AUTHOR (S):

Scalia, Rosario; Gefen, Jonathan; Petasis, Nicos A.;

Serhan, Charles N.; Lefer, Allan M.

CORPORATE SOURCE:

Dep. Physiology, Jefferson Medical College, Thomas Jefferson Univ., Philadelphia, PA, 19107-6799, USA

SOURCE:

Proc. Natl. Acad. Sci. U. S. A. (1997), 94(18),

9967-9972

CODEN: PNASA6; ISSN: 0027-8424 National Academy of Sciences

DOCUMENT TYPE:

Journal

PUBLISHER: LANGUAGE:

English

Three different stable lipoxin A4 (LXA4) analogs (i.e., AB 16-phenoxy-LXA4-Me, 15-cyclohexyl-LXA4-Me, and 15-R/S-methyl-LXA4-Me) were studied of their ability to modulate leukocyte-endothelial cell interactions in the rat mesenteric microvasculature. Superfusion of the rat mesentery with 50 .mu.mol/L NG-nitro-L-arginine Me ester (L-NAME) caused a significant, time-dependent increase in leukocyte rolling (56.+-.8 cells/min; vs. control) and leukocyte adherence (12.5.+-.1.2 Searched by Barb O'Bryen, STIC 308-4291

cells/100 .mu.m length of venule; vs. control) after 120 min of superfusion. Concomitant superfusion of the rat mesentery with 10 nmol/L of each of three lipoxin analogs consistently and markedly attenuated L-NAME-induced leukocyte rolling to 10.+-.4, 4.+-.1, and 32.+-.7 cells/min, and adherence to 4.+-.0.8, 1.1.+-.0.4, and 7.+-.0.7 cells/100 .mu.m length of venule (16-phenoxy-LXA4-Me, 15-cyclohexyl-LXA4-Me, and 15-R/S-methyl-LXA4-Me, resp.). No alterations of systemic blood pressure or mesenteric venular shear rates were obsd. in any group. Immunohistochem. up-regulation of P-selectin expression on intestinal venular endothelium was significantly increased after exposure to L-NAME, and this was significantly attenuated by these lipoxin analogs. Thus, in vivo superfusion of the rat mesentery with stable lipoxin analogs at 10 nmol/L reduces L-NAME-induced leukocyte rolling and adherence in the mesenteric rat microvasculature by attenuating P-selectin expression. This anti-inflammatory mechanism may represent a novel and potent regulatory action of lipoxins on the immune system.

IT 171030-14-1

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses)

(lipoxin A4 stable analogs inhibit leukocyte rolling and adherence in rat mesenteric microvasculature)

171030-14-1 CAPLUS RN

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, methyl CNester, (5S, 6R, 7E, 9E, 11Z, 13E, 15R) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

L12 ANSWER 19 OF 23 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1997:196043 CAPLUS

DOCUMENT NUMBER:

126:291403

TITLE:

Lipoxin A4 stable analogs are potent mimetics that

stimulate human monocytes and THP-1 cells via a

G-protein-linked lipoxin A4 receptor

AUTHOR(S):

Maddox, Jane F.; Hachicha, Mohamed; Takano, Tomoko; Petasis, Nicos A.; Fokin, Valery V.; Serhan, Charles

CORPORATE SOURCE:

Cent. Exp. Therapeut. Reperfusion Injury, Brigham and

Women's Hosp. and Harvard Med. Sch., Boston, MA,

02115, USA

SOURCE:

J. Biol. Chem. (1997), 272(11), 6972-6978

CODEN: JBCHA3; ISSN: 0021-9258

PUBLISHER:

American Society for Biochemistry and Molecular

Biology

DOCUMENT TYPE:

Journal English

LANGUAGE:

Lipoxins (LX) are bioactive eicosanoids that activate human monocytes and inhibit neutrophils. LXA4 is rapidly converted by monocytes to inactivate products, and to resist metab., synthetic analogs of LXA4 were designed. Here, the authors examd. the bioactivity of several LXA4 analogs in monocytes and found, for chemotaxis, 15(R/S)-methyl-LXA4 were equal in activity, and 16-phenoxy-LXA4 was more potent than native LXA4. 15(R/S)-methyl-LXA4 and 16-phenoxy-LXA4 were .apprx.1 log molar more Searched by Barb O'Bryen, STIC 308-4291

potent than LXA4 in stimulating THP-1 cell adherence (EC50 .apprxeq.1 .times. 10-10 M). Dimethylamide derivs. of the LXA4 analogs also possessed agonist rather than antagonist properties for monocytes. Neither LXA4 nor 16-phenoxy-LXA4 affected monocyte-mediated cytotoxicity. The authors cloned an LXA4 receptor from THP-1 cells identical to that found in PMN. Evidence of receptor-mediated function of LXA4 and the stable analogs in monocytes included desensitization of intracellular calcium mobilization to a second challenge by equimolar concns. of these analogs, but not to LTB4. Increases in [Ca2+]i by LXA4 and the analogs were specifically inhibited by an antipeptide antibody to the LXA4 receptor; and both LXA4- and analog-induced adherence and increments in Ca2+ were sensitive to pertussis toxin. Together, these results indicate that the LXA4 stable analogs are potent monocyte chemoattractants and are more potent than native LXA4 in stimulating THP-1 cell adherence, at subnanomolar concns. Moreover, they provide addnl. evidence that the LXA4 stable analogs retain selective bioactivity in monocytes and are valuable instruments for examg. the functions and modes of action of LXA4.

IT 171030-12-9 189005-35-4

RL: BAC (Biological activity or effector, except adverse); PRP (Properties); BIOL (Biological study)

(lipoxin A4 stable analogs are potent mimetics that stimulate human monocytes and THP-1 cells via a G-protein-linked lipoxin A4 receptor)

RN 171030-12-9 CAPLUS

CN 7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, (5s,6R,7E,9E,11Z,13E,15R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

RN 189005-35-4 CAPLUS

CN 7,9,11,13-Hexadecatetraenamide, 5,6,15-trihydroxy-N,N-dimethyl-16-phenoxy-, [5S-(5R*,6S*,7E,9E,11Z,13E,15S*)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.
Double bond geometry as shown.

$$Me_2N$$
 OH E E E E OPh OH

L12 ANSWER 20 OF 23 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1995:435841 CAPLUS

DOCUMENT NUMBER: 122:205187

TITLE: Lipoxin compounds for modulation of inflammation of

columnar epithelia

INVENTOR(S): Madara, James L.; Serhan, Charles N.; Colgan, Sean P.

PATENT ASSIGNEE(S): Brigham and Women's Hospital, USA

SOURCE: PCT Int. Appl., 60 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English '

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

APPLICATION NO. DATE PATENT NO. KIND WO 1994-US7333 WO 9501179 **A**1 19950112 AT, AU, BB, BG, BR, BY, CA, CH, CN, CZ, DE, DK, ES, FI, GB, HU, JP, KP, KR, KZ, LK, LU, LV, MG, MN, MW, NL, NO, NZ, PL, PT, RO, RU, SD, SE, SK, UA, UZ, VN RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE AU 1994-72152 19940629 19950124 AU 9472152 **A1** US 1993-84311 19930629 PRIORITY APPLN. INFO.: 19940629 WO 1994-US7333

MARPAT 122:205187 OTHER SOURCE(S):

A pharmaceutical compn. for treating or preventing a disease or condition assocd. with columnar epithelial inflammation or with abnormal transportation of fluids, electrolytes, or nutrients by a columnar epithelium contains lipoxin A4 or its analogs. Columnar epithelium is an epithelium of the intestine, kidney, stomach, liver, thyroid, trachea, lung, gall bladder, urinary bladder, bile duct, pancreatic duct, or testicle.

161718-15-6 161718-22-5 IT

RL: BAC (Biological activity or effector, except adverse); THU (Therapeutic use); BIOL (Biological study); USES (Uses) (lipoxin compds. as inflammation inhibitors for columnar epithelia)

__161718-15-6 CAPLUS RN

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy- (9CI) CN (CA INDEX NAME)

PAGE 1-A OH OH OH

PAGE 1-B

CO2H

RN 161718-22-5 CAPLUS 7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, methyl CN (CA INDEX NAME) ester (9CI)

PAGE 1-A OH он он

PAGE 1-B

OPh

L12 ANSWER 21 OF 23 CAPLUS COPYRIGHT 2001 ACS

1995:875227 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 124:213

Design of Lipoxin A4 Stable Analogs That Block TITLE:

Transmigration and Adhesion of Human Neutrophils Serhan, Charles N.; Maddox, Jane F.; Petasis, Nicos

AUTHOR (S): A.; Akritopoulou-Zanze, Irini; Papayianni, Aikaterina;

Brady, Hugh R.; Colgan, Sean P.; Madara, James L. Center for Experimental Therapeutics and Reperfusion

Injury, Brigham and Women's Hospital, Boston, MA,

02115, USA

Biochemistry (1995), 34(44), 14609-15 SOURCE:

CODEN: BICHAW; ISSN: 0006-2960

DOCUMENT TYPE:

CORPORATE SOURCE:

Journal LANGUAGE: English

Lipoxins (LX) are bioactive eicosanoids that carry a tetraene structure and serve as regulators of inflammation, in part by inhibiting neutrophil migration and adhesion. Lipoxin A4 is rapidly regulated by conversion to inactive LX metabolites via local metab. that involves dehydrogenation as the predominant route. Here, several LXA4 analogs were designed that resisted rapid conversion by both differentiated HL-60 cells and recombinant 15-hydroxyprostaglandin dehydrogenase, systems where native LXA4 is degraded within minutes. The rank order of conversion by recombinant dehydrogenase was LXA4 Me ester > PGE2 .apprxeq. PGE2 Me ester > LXA4 >>> the novel LXA4 analogs. In addm., 15(R/S)-methyl-LXA4, 15-cyclohexyl-LXA4, and 16-phenoxy-LXA4 proved to retain LXA4 bioactivity and inhibited neutrophil transmigration across polarized epithelial cell monolayers as well as adhesion to vascular endothelial cells. These results indicate that LXA4 analogs can be designed using these criteria to resist rapid transformation and to retain biol. actions of native LXA4. Moreover, the results suggest that LXA4 stable analogs can be useful tools both in vitro and in vivo to evaluate LXA4 actions and therapeutic potential.

IT 171030-12-9

RL: BAC (Biological activity or effector, except adverse); BPR (Biological process); BIOL (Biological study); PROC (Process)

(design of lipoxin A4 stable analogs that block transmigration and adhesion of human neutrophils)

RN171030-12-9 CAPLUS

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, CN (5S, 6R, 7E, 9E, 11Z, 13E, 15R) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

IT 171030-14-1

RL: BPR (Biological process); BIOL (Biological study); PROC (Process) (design of lipoxin A4 stable analogs that block transmigration and adhesion of human neutrophils)

171030-14-1 CAPLUS RN

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, methyl CN ester, (5S, 6R, 7E, 9E, 11Z, 13E, 15R) - (9CI) (CA INDEX NAME)

Absolute stereochemistry. Double bond geometry as shown.

L12 ANSWER 22 OF 23 CAPLUS COPYRIGHT 2001 ACS

1995:573821 CAPLUS ACCESSION NUMBER:

122:314353 DOCUMENT NUMBER:

TITLE: Lipoxin compounds INVENTOR (S): Serhan, Charles N.

Brigham and Women's Hospital, USA PATENT ASSIGNEE(S):

PCT Int. Appl., 99 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

	PATENT NO.	KIND DATE	APPLICATION NO. DATE
	WO 9429262	A1 19941222	WO 1994-US6822 19940615
	W: AT, AU	, BB, BG, BR, BY,	CA, CH, CN, CZ, DE, DK, ES, FI, GB, HU,
	JP, KP	, KR, KZ, LK, LU,	LV, MG, MN, MW, NL, NO, NZ, PL, PT, RO,
		, SE, SK, UA, UZ,	v v v v v v v v v v v v v v v v v v v
	RW: AT, BE	, CH, DE, DK, ES,	FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
	CA 2164951	AA 19941222	CA 1994-2164951 19940615
	AU 9471109	A1 19950103	AU 1994-71109 19940615
	AU 692453	B2 19980611	
			EP 1994-920241 19940615
	R: AT, BE	, CH, DE, DK, ES,	FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE
	JP 08512023	T2 19961217	JP 1994-502216 19940615
PRI	ORITY APPLN. INFO	o.;	US 1993-77300 19930615
			WO 1994-US6822 19940615
ОТН	ER SOURCE(S)	MARPAT 122:	314353

OTHER SOURCE(S):

GΙ

Compds. having the active site of natural lipoxins, but a longer tissue Searched by Barb O'Bryen, STIC 308-4291 AB

half-life, in particular I [R = H, Me; R1 = pentyl, cyclohexyl, CH2OPh] and their 11,12-didehydro analogs are disclosed. These small mols. are useful for treating vasoconstrictive, inflammatory, myeloid suppressive, cardiovascular, and gastrointestinal diseases. Thus, I inhibited neutrophil adhesion to endothelial cells and polymorphonuclear cell adhesion to endothelial cells triggered by leukotriene B4. 1He acetylenic analogs of I were more stable than I.

IT 161718-15-6P 161718-22-5P

RL: BAC (Biological activity or effector, except adverse); IMF (Industrial manufacture); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(lipoxin analogs with longer tissue half-life)

RN 161718-15-6 CAPLUS

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy- (9CI) (CA INDEX NAME)

PAGE 1-B

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CN

RN 161718-22-5 CAPLUS CN 7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, methyl ester (9CI) (CA INDEX NAME)

PAGE 1-B

— oph

L12 ANSWER 23 OF 23 USPATFULL

ACCESSION NUMBER:

97:61839 USPATFULL

TITLE:

Lipoxin compounds

INVENTOR(S):

PATENT ASSIGNEE(S):

Serhan, Charles N., Boston, MA, United States

Brigham & Womens Hospital, Boston, MA, United States

(U.S. corporation)

NUMBER

DATE

PATENT INFORMATION:

US 5648512 19970715

APPLICATION INFO .:

US 1995-453125 19950531

RELATED APPLN. INFO.:

Division of Ser. No. US 1994-260030, filed on 15 Jun 1994, now patented, Pat. No. US 5441951 which is a continuation-in-part of Ser. No. US 1993-77300, filed

on 15 Jun 1993, now abandoned

DOCUMENT TYPE:

Utility Geist, Gary

PRIMARY EXAMINER: ASSISTANT EXAMINER:

Williams, Rosalynd

LEGAL REPRESENTATIVE:

Lahive & Cockfield, LLP

NUMBER OF CLAIMS: EXEMPLARY CLAIM:

1

LINE COUNT:

2197

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB

Compounds having the active site of natural lipoxins, but a longer tissue hlf-life are disclosed. These small molecules are useful for treating vasoconstrictive, inflammatory, myeloid suppressive, cardiovascular, and gastrointestinal diseases.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

161718-15-6P 161718-22-5P

(lipoxin analogs with longer tissue half-life)

161718-15-6 USPATFULL RN

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy- (9CI) CN (CA INDEX NAME)

PAGE 1-A он он OH PhO-CH2-CH-CH-CH-CH-CH-CH-CH-CH-CH-CH-CH-CH-(CH2)3-

PAGE 1-B

CO2H

161718-22-5 USPATFULL RN

7,9,11,13-Hexadecatetraenoic acid, 5,6,15-trihydroxy-16-phenoxy-, methyl CN ester (9CI) (CA INDEX NAME)

PAGE 1-A OH ' 0 OHOH

PAGE 1-B

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